Scheduling non-identical jobs on a batch resource

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Introduction

This paper discusses three different approaches, and several variations on them, to solving the problem of scheduling non-identical jobs on a batch processing machine. Batch processing machines, for the purposes of this paper, can process multiple, non-identical jobs simultaneously—but all jobs must be loaded into and unloaded from the machine at once, which introduces a considerable twist on the "simple parallel resources" known from typical example problems in existing literature.

The machines in question represents real-life resources like autoclaves or ovens, which can process multiple items at a time, but often cannot be opened at random—in fact, such machines often need to wait for the largest item in the batch to be done before the next batch can be inserted.

Malapert et al. [2012] proposed a global constraint programming algorithm consisting of a set of filtering rules to solve the problem. He achieved considerably better speeds than with a simple mixed-integer model, but it seems plausible that the new global constraint is unnecessarily cumbersome to achieve this performance; simple MIP, CP or decomposition approaches are easier to implement and extend. In this paper, we present 1) an improvement to his MIP model, 2) a CP model and 3) a decomposition approach to "divide and conquer" the problem.

1.1 Problem definition

We describe the problem as follows: assume we are given a set of jobs J, each of which has a processing time (or "length") p_j and a size (or "capacity requirement") s_j . Each job also has a due date d_j . The machine is characterized by its capacity b, and in every batch, the jobs' summed sizes must not exceed this number. All values are integer.

The machine can only process one batch k of jobs at a time, and batches always take as long as the longest job in the batch (i.e. $P_k = \max_{j \in k}(p_j)$). Our objective is to minimize the lateness L of the latest job in J, where L is the difference between the job's completion time C_j and its due date d_j —in formal terms, min. $L_{\max} = \max_j (C_j - d_j)$. The job's completion time, however, is the completion time of the batch, which in turn finishes with its *longest* job as stated above.

Malapert uses the standard format established by Graham et al. to summarize the problem as 1|p-batch;b| < n; non-identical $|L_{\max}|$, where p-batch;b| < n represents the parallel-batch nature and the finite capacity of the resource. A simpler version with identical job sizes was shown to be strongly NP-hard in [Brucker et al., 1998]; this problem, then, is no less difficult.

It helps to visualize the jobs before delving into the technicalities of scheduling them. Figure 1.1 shows a solution to a sample problem with eight jobs and a resource with capacity b = 20.

1.2 Organization of this paper

After reviewing some of the most relevant publications on both general MIP/CP models and batch scheduling problems, we first describe Malapert's original MIP model in section 3.1. We then present possible improvements to the model in 3.2. Section 3.3 introduces a CP formulation of the same problem. Sections 3.4.1 and 3.4.1 describe a decomposition approach.

An empirical comparison of the new models and a discussion of the results follow in

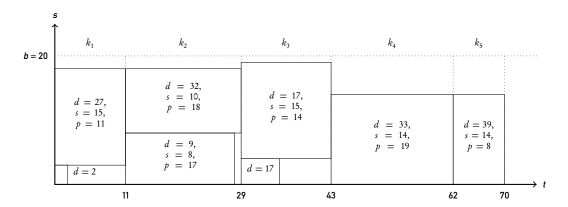


Figure 1.1: Optimal solution to an example problem with eight jobs (s_j and p_j not shown for the two small jobs)

sections 4.1 and 5. Ideas for future work are listed in 6.

Background

Many optimization problems, and scheduling problems in particular, are combinatorial in nature. The number of possible solutions grows exponentially with the number of input variables, and even with fast computers it is impossible to explore all of them individually to find the best one (also called "full enumeration", or "brute-force" search) in a reasonable amount of time. Often, however, it is possible to reason about subsets of solutions that are known to be suboptimal a priori. This limits the search space, allowing us to solve many instances of difficult combinatorial problems in few hours, minutes or even seconds.

Such constrained searches are often implemented in either of two ways (or variants of them): as a *Constraint Programming model* (CP) or as a *Mixed Integer Programming model* (MIP). In this section I will briefly introduce the concepts behind both CP and MIP and review recent work on problems similar to the one dealt with here.

2.1 Constraint Programming

A constraint program specifies relationships, *constraints*, that must hold between input variables in the form of mathematical equations or inequalities. Every input variable is assigned a set \mathcal{D} of potential values it can assume, its *domain*. The solving algorithm begins by assigning a value to one of the variables. Using the constraints, the solver can then often reduce the other variables' domains, i.e. eliminate other potential values of other variables. These domain reductions may themselves trigger domain reductions in other variables, and so on, in a process called *propagation*. Propagation can lead to one of three results:

- an inconsistency, in which a variable's domain is reduced to an empty set. That is, the given constraints allow for none of that variable's potential values given the values assigned to other variables. As a result, the solver will *backtrack*, i.e. undo one or more of previous assignments made, and assign other values.
- a consistent solution, in which every variable's domain is reduced to a single value (a singleton). If we are searching for an optimal solution, this may replace an incumbent solution if it is better.
- some reduced domains, but neither a solution nor an inconsistency. In this case, the solver assigns more values to variables to trigger more propagation, until a solution or an inconsistency is reached.

Without formally introducing different notions of inconsistency, it should be noted that a problem can have no solution even when all variables have non-empty domains. (See figure).

Unlike in MIP, constraints in CP are not restricted to expressions that are linear in the input variables. Furthermore, so-called *global constraints* can be used that act on entire sets of variables at once. They are implemented as custom algorithms that rely on specific knowledge about the problem and are often based on results from graph theory. Global constraints can be powerful tools; a classic example is the alldifferent constraint which ensures that a set of variables take on unique values. While a number of not-equal relationships between the variables can achieve the same effect after several propagations, the global constraint reduces the domains of all involved variables in one fell swoop such that for each variable, only the values remain that are consistent with those of all the other variables.

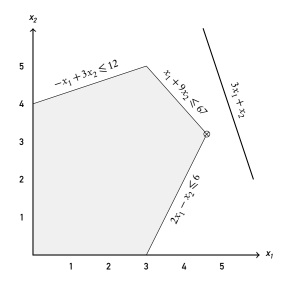


Figure 2.1: Graphical representation of a typical LP problem

2.2 Mixed Integer Programming

2.2.1 Linear Programming

Mixed Integer Programs (or "MIP models") express the minimization of a linear function subject to linear constraints. If all variables in the problem can be rational in the solution, the MIP model is really a *linear program* (LP), which can be solved in polynomial time.¹

Figure 2.1 illustrates the concept of an LP in two variables, x_1 and x_2 , as listed in model 2.2.1. The set of feasible solutions is given by the shaded area bounded by the axes and by three inequalities ("constraints"). A third linear term, the objective function $3x_1 + x_2$, is to be maximized.²

Although the objective function is shown in the figure as a line in a specific location, note that, as it is not an equation, it can be represented by any line parallel to that shown in the figure. While we are trying to maximize the objective value, the solution must be feasible. It is thus obvious that the desired extreme value of our objective function is found

¹In practice, variations on Dantzig's *simplex method* are most often used to solve LPs. Such solvers perform very well on most problems, but no known variant has been proven to have polynomial worst-case complexity [?]. Solvers with theoretically polynomial-time complexity exist (Karmarkar's algorithm? has a runtime of $\mathcal{O}(n^{3.5}L^2 \cdot \log L \cdot \log \log L)$, for instance, where L is the number of bits of input), but are used less frequently.

²Minimization is more common, but note that multiplying the objective by -1 achieves this.

Maximize
$$3x_1 + x_2$$
 (2.1)

subject to the constraints
$$-x_1 + 3x_2 \le 12$$
 (2.2)

$$x_1 + 9x_2 \le 67 \tag{2.3}$$

$$2x_1 - x_2 \leqslant 6 \tag{2.4}$$

$$x_1 \geqslant 0 \tag{2.5}$$

$$x_2 \geqslant 0 \tag{2.6}$$

Model 2.2.1: A simple LP model, as shown in figure 2.1

at one of the "corners" of the shaded area: the solution is marked \otimes in the figure. In fact, since the shaded area generated by linear inequalities will always be a convex polygon (or polyhedron, in higher dimensions), the solution will invariably be found at an intersection of hyperplanes.

2.2.2 Solving MIP models using branch-and-bound

MIP models are LPs in which some of the decision variables are declared integers—thus their name. Like LP models, MIP models also require an objection function to be minimized. Figure 2.3 illustrates this based on the LP problem above: now, only the black dots represent feasible solutions. While the solution is easily found in the figure by inspection (simply round to the nearest feasible integral solution!), this is not the case in problems with many variables; it is difficult enough to visualize the problem in three dimensions, and many problems require hundreds or thousands. Moreover, rounding is particuarly unreliable with variables of small domains (e.g. binary variables), which are often used in MIP models to represent decisions. Indeed, solving MIP models is NP-hard.

Similar to CP searches, MIP searches can be thought of as trees, where every branch represents an assignment of a value to a domain and every leaf represents a feasible solution. The difference lies in the way MIP explores this search tree.

The MIP solver first solves the problem as an LP, which will usually result in fractional values for all or most of the variables. In this context, the LP is known as the *LP relaxation*

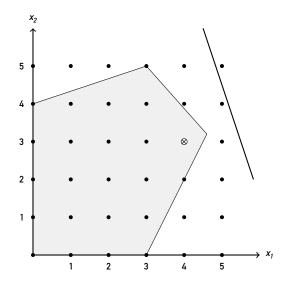


Figure 2.2: Graphical representation of a typical MIP problem

of the problem—an easier, approximate version of the original. Assuming we are minimizing the objective, the resulting LP objective value will be a lower bound on the optimal MIP objective value. At this point, the solution is $x_1 = 4.6$, $x_2 = 3.2$.

The solver then chooses a variable, based on heuristics, and *branches* on it. In this case, assume that we branch on x_2 , which means that we set $x_2 \le 3.0 \lor x_2 \le 4.0$. We now have two new MIPs, as shown in figure ??.

. The current solution from the LP relaxation is fractional and thus infeasible. The search tree

which generally results in fractional values for most variables, such as x = a. It then branches on possible values for x: either x < a or x > a. If a is integral, x = a. The MIP solution at a node n also serves as a lower bound LB_n on solutions in the respective subtrees: once a solution with an objective value of v is found, v becomes an upper bound on the optimal objective value and all yet unexplored subtrees with $LB_n \ge v$ need no longer be searched ("pruning" the subtree). This procedure is called branch-and-bound and the most basic method used to solve integer programming problems. Other methods exist, e.g. branch-and-cut, and are often used in conjunction with branch-and-bound in today's solvers.

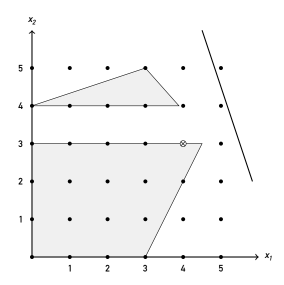


Figure 2.3: Graphical representation of a typical MIP problem

2.3 Literature

The problem at hand is based on the work of Malapert et al. [2012], who proposed a global constraint sequenceEDD to be used in combination with pack to solve it to optimality. The sequenceEDD constraint is implemented as four distinct filtering rules applied at relevant domain changes: three to update bounds on $L_{\rm max}$ based on different conditions, and one to limit the number of batches based on the marginal cost difference between adding a job to an empty batch vs. an existing one. The new global was shown to significantly outperform a simple MIP model of the same problem.

Other authors have examined similar problems: Azizoglu and Webster [2000] provide an exact method and a heuristic for the same problem, but minimize makespan (C_{max}) instead of L_{max} , as have Dupont and Dhaenens-Flipo [2002]. Similar exact methods have been proposed for multi-agent variants with different objective functions [Sabouni and Jolai, 2010], for makespan minimization on single batch machines [Kashan et al., 2009], and for makespan minimization on parallel batch machines with different release dates [Ozturk et al., 2012]. A more extensive review of MIP model applications in batch processing is given by Grossmann [1992].

Modelling the problem

3.1 MIP model

As a baseline to compare other models' performance to, we first replicated Malapert's original MIP approach, as given in Model 6.5.1. It uses a set of binary decision variables x_{jk} to represent whether job j is assigned to batch k. The original model assumes a set K of |K| = |J| batches, the number of jobs being a trivial upper bound on the number of batches required; it also enforces an earliest-due-date-first (EDD) ordering of the batches (constraint 3.7).

Min.
$$L_{\text{max}}$$
 (3.1)

s.t.
$$\sum_{k \in K} x_{jk} = 1 \qquad \forall j \in J$$
 (3.2)

$$\sum_{j \in J} s_j x_{jk} \leqslant b \qquad \forall k \in K \tag{3.3}$$

$$p_j x_{jk} \leqslant P_k \qquad \qquad \forall j \in J, \forall k \in K \tag{3.4}$$

$$C_{k-1} + P_k = C_k \qquad \forall k \in K \tag{3.5}$$

$$(d_{max} - d_j)(1 - x_{jk}) + d_j \geqslant D_k \quad \forall j \in J, \forall k \in K$$

$$(3.6)$$

$$D_{k-1} \leqslant D_k \qquad \qquad \forall k \in K \tag{3.7}$$

$$C_k - D_k \leqslant L_{\text{max}} \qquad \forall k \in K \tag{3.8}$$

$$C_k \geqslant 0, P_k \geqslant 0 \text{ and } D_k \geqslant 0 \qquad \forall k \in K$$
 (3.9)

Model 3.1.1: Malapert's original MIP model

Improved MIP model 3.2

Several improvements can be made to Malapert's MIP model in the form of additional constraints shown in Model 3.2.1, as described in greater detail in the subsections below.

$$\sum_{j \in I} x_{j,k-1} = 0 \to \sum_{j \in I} x_{jk} = 0 \quad \forall k \in K$$

$$(3.10)$$

$$e_k + \sum_{i \in I} x_{jk} \geqslant 1 \qquad \forall k \in K \tag{3.11}$$

$$n_{j}(e_{k}-1)+\sum_{j\in I}x_{jk}\leqslant 0 \qquad \forall k\in K \tag{3.12}$$

$$e_k - e_{k-1} \geqslant 0 \qquad \forall k \in K \tag{3.13}$$

$$x_{jk} = 0 \qquad \qquad \forall \{j \in J, k \in K | j > k\} \tag{3.14}$$

$$\begin{aligned} e_k - e_{k-1} &\geqslant 0 & \forall k \in K & (3.13) \\ x_{jk} &= 0 & \forall \{j \in J, k \in K | j > k\} & (3.14) \\ L_{\max} &\geqslant \left\lceil \frac{1}{b} \sum_j s_j p_j \right\rceil - \delta_q & \forall q, \forall \{j \in J | d_j \leqslant \delta_q\} & (3.15) \end{aligned}$$

Model 3.2.1: Improvements to Malapert's original MIP model

3.2.1 Grouping empty batches

The given formulation lacks a rule that ensures that no empty batch is followed by a nonempty batch. Empty batches have no processing time and a due date only bounded by d_{max} , so they can be sequenced between non-empty batches without negatively affecting L_{\max} . Since, however, desirable schedules have no empty batches scattered throughout, we can easily reduce the search space by disallowing such arrangements. The idea is illustrated in Figure 3.1.

A mathematical formulation is

$$\sum_{j \in J} x_{j,k-1} = 0 \to \sum_{j \in J} x_{jk} = 0 \quad \forall k \in K.$$
 (3.16)

To implement this, we can write constraints in terms of an additional set of binary variables, e_k , indicating whether a batch k is empty or not:

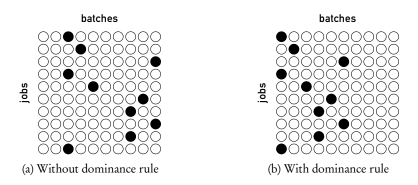


Figure 3.1: Dominance rule to eliminate empty batches followed by non-empty batches (circles represent the x_{jk} variables; a filled circle stands for $x_{jk} = 1$)

$$e_k + \sum_{j \in I} x_{jk} \geqslant 1 \qquad \forall k \in K, \tag{3.17}$$

$$n_j(e_k - 1) + \sum_{j \in I} x_{jk} \leqslant 0 \quad \forall k \in K.$$
 (3.18)

Constraints (3.17) enforce $e_k = 1$ when the batch k is empty. Constraints (3.18) enforce $e_k = 0$ otherwise, since the sum term will never exceed n_j . The rule (3.16) can now be expressed as $e_{k-1} = 1 \rightarrow e_k = 1$, and implemented as follows:

$$e_k - e_{k-1} \geqslant 0 \quad \forall k \in K. \tag{3.19}$$

We can also prune any attempts to leave the first batch empty by adding a constraint $e_0 = 0$.

3.2.2 No postponing of jobs to later batches

Since the jobs are already sorted by non-decreasing due dates, it makes sense to explicitly instruct the solver never to attempt to push jobs into batches with a greater index than their own: even if every job had its own batch, it would be unreasonable to ever postpone

a job to a later batch.

$$x_{jk} = 0 \quad \forall \{j \in J, k \in K | j > k\}$$
 (3.20)

3.2.3 Lower bound on L_{max}

Let a bucket q denote the set of all batches with due date δ_q . Then the completion date C_q of this bucket is the completion date of the last-scheduled batch with due date δ_q , and the lateness of the bucket q is $L_q = C_q - \delta_q$. Since all batches up to and including those in bucket q are guaranteed to contain all jobs with due dates $d \leq \delta_q$ – as ensured by the EDD ordering of batches – the lower bound on every bucket's lateness $LB(L_q)$ is a valid lower bound on L_{\max} . In other words, jobs with due date $d \leq \delta_q$ will be found only in batches up to and including the last batch of bucket q. This provides a lower bound on the lateness of bucket q:

$$L_{\max} \geqslant C_{\max,q} - \delta_q \quad \forall q \tag{3.21}$$

The buckets up to bucket q will likely also contain some later $(d > \delta_q)$ jobs in the optimal solution but this does not affect the validity of the lower bound.

Now we need to find $C_{\max,q}$, or at least a lower bound on it, in polynomial time. The simplest approach simply considers the jobs' total "area" (or "energy"), i.e. the sum of all $s_i p_i$ products:

$$C_{\max,q} \geqslant \lceil \frac{1}{b} \sum_{j} s_{j} p_{j} \rceil \quad \forall q, \forall \{j \in J | d_{j} \leqslant \delta_{q} \}$$
 (3.22)

A better lower bound on $C_{\max,q}$ would be given by a preemptive-cumulative schedule. Unfortunately, minimizing C_{\max} for such problems is equivalent to solving a standard bin-packing problem, which requires exponential time.¹

¹In a preemptive-cumulative schedule, jobs may be stopped and restarted mid-execution, but occupy a constant amount s_j on the resource while executing. In such a schedule, minimizing the makespan is as difficult as solving a bin-packing problem: we can break jobs into small pieces (no longer than the smallest common divisor of the jobs' lengths p) and then pack them together such as to minimize the number of small time slots needed.

3.3 CP model

The constraint programming model is based on a set of decision variables $B_j = \{k_1, ..., k_{n_k}\}$, where each variable stands for the batch job j is assigned to. The complete model is given in Model 3.3.1.

Min.
$$L_{\text{max}}$$
 (3.23)

s.t.
$$\operatorname{pack}(J, K, b)$$
 (3.24)

$$cumul(J,b) (3.25)$$

$$P_k = \max_j p_j \qquad \qquad \forall \{j \in J | B_j = k\}, \forall k \in K$$
 (3.26)

$$D_k = \min_{j} d_j \qquad \qquad \forall \{j \in J | B_j = k\}, \forall k \in K$$
 (3.27)

$$C_k + P_{k+1} = C_{k+1} \qquad \forall k \in K \tag{3.28}$$

$$L_{\max} \geqslant \max_{k} \left(C_k - D_k \right) \tag{3.29}$$

$$B_{j} \leq k \qquad \qquad \forall \{j \in J, k \in K | j > k\} \tag{3.31}$$

Model 3.3.1: Constraint programming model

Constraint (3.24) makes sure the jobs are distributed into the batches such that no batch exceeds the capacity b. Constraint (3.25), a global cumulative constraint, keeps jobs from overlapping on the resource – while redundant with (3.24), this speeds up propagation slightly. Constraints (3.26), (3.27), (3.28) and (3.29) define P_k , D_k , C_k (batch completion time) and $L_{\rm max}$, respectively.

3.3.1 Grouping empty batches

We can force empty batches to the back and thus establish dominance of certain solutions. The implementation is much easier than in the MIP model:

3.3.2 No postponing of jobs to later batches

Just like in the MIP model, jobs should never go into a batch with an index greater than their own:

$$B_i \le k \quad \forall \{j \in J, k \in K | j > k\} \tag{3.33}$$

3.4 Decomposition approach

Instead of solving the entire problem using one model, we can solve the problem step by step, using the best techiques available for each subproblem. This approach is inspired by a method called *Benders decomposition*.

3.4.1 Branch-and-bound by batch in chronological order

A basic version of this approach uses branch-and-bound to transverse the search tree. At each node on level ℓ , a single MIP and/or CP model is run to assign jobs to batch $k=\ell$. The remaining jobs are passed to the children nodes, which assign jobs to the next batch, and so on – until a solution, and thus a new upper bound on $L_{\rm max}$, is found. Several constraints are used to prune parts of the search tree that are known to offer only solutions worse than this upper bound. Figure 3.2 shows an example in which a MIP model is used to assign jobs to the batch at every node. Algorithm 1 outlines what happens at each node: the model finds the best jobs to assign to the batch according to some rule, lets the children handle the remaining jobs, and tries the next best solution once the first child has explored its subtree and backtracked.

Using MIP and cumulative packing after the batch

The first version of the branch-and-bound batch-by-batch decomposition uses a MIP model at each node to assign jobs to the respective batch. The remaining jobs are packed such as to minimize their $L_{\rm max}$, with a relaxation of the batching requirement, i.e., as if on a cumulative resource.

Algorithm 1 MIP node class code overview

```
update currentAssignments
                                                                                    b this keeps track of where we are in the tree
if no jobs given to this node then
                                                                       ▶ if this is a leaf node, i.e. all jobs are assigned to batches
    calculate L_{\text{max,current}} based on currentAssignments
    if L_{\text{max,current}} < L_{\text{max,incumbent}} then
        L_{\text{max,incumbent}} \leftarrow L_{\text{max,current}}
        bestAssignments \leftarrow currentAssignments
    end if
    return to parent node
end if
set up MIP model
                                                                                                                > as described below
repeat
                                                                                               ▶ let model assign jobs to the batch
    x_i \leftarrow \text{model.solve}(x_i)
    spawn and run child node with all \{j | x_i = 0\}
                                                                                                 ▶ pass unassigned jobs to children
    add constraint to keep this solution from recurring
                                                                                      ▶ this happens once the child node returns
until model has no more solutions
return to parent node
```

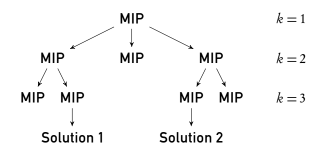


Figure 3.2: Batch-by-batch decomposition using MIP only

Model 3.4.1 implements a time-indexed cumulative constraint on the non-batched jobs. Constraints (3.35) through (3.37) ensure that the batch stays below capacity, define the duration of the batch P_k and force at least one of the earliest-due jobs into the batch.

Constraints (3.38) express the interval relaxation used to ensure that no jobs exceed their latest allowable finish date given any batch assignment. Even jobs that are assigned to the batch have to fulfill this requirement.

Constraints (3.39) and (3.40) implement the cumulative nature of the post-batch assignments by ensuring that each job starts only once, and no jobs overlap on a given resource at any time.

Constraints (3.41) again limit the possible end dates of a job, but unlike (3.38), they use the time assignments on the cumulative resource to determine end dates. Constraints (3.42)

Min.
$$L_{\text{max,cumul}}$$
 (3.34)

s. t.
$$\sum_{j} s_{j} x_{j} \leq b$$
 $\forall j \in J$ (3.35)

$$P_k \geqslant p_j x_j \qquad \forall j \in J \tag{3.36}$$

$$\sum_{i} x_{j} \geqslant 1 \qquad \qquad \forall \{j \in J | d_{j} = \min(d_{j})\} \tag{3.37}$$

$$P_k + \frac{1}{b} \sum_{i} s_i p_i \leqslant d_j + L_{\text{max,incmb}} - 1 - v_k \quad \forall j \in J, \forall \{i \in J | d_i \leqslant d_j\}$$
 (3.38)

$$\sum u_{jt} = 1 \qquad \forall j \in J \tag{3.39}$$

$$\sum_{j} \sum_{t' \in T_{i,t}} s_j u_{jt'} \leqslant b \qquad \forall t \in \mathcal{H}$$
 (3.40)

$$(v_k + t + p_j)u_{jt} \le d_j + L_{\text{max.incmb}} - 1 \qquad \forall j \in J, \forall t \in \mathcal{H}$$
(3.41)

$$L_{\text{max,cumul}} \ge (v_k + t + p_i)u_{it} - d_i \qquad \forall j \in J, \forall t \in \mathcal{H}$$
 (3.42)

$$u_{j,t=0} = x_j \qquad \forall j \in J \tag{3.43}$$

$$u_{it} \le (1 - x_j)$$
 $\forall i, j \in J, \forall t \in \{1, ..., p_j - 1\}$ (3.44)

$$b - \sum_{i \in I} s_i x_i \le (b w_j + 1) s_j \qquad \forall j \in J$$
 (3.45)

$$P_k + 2w_j n_t \geqslant p_j + n_t x_j \qquad \forall j \in J$$
 (3.46)

$$P_k - 2(1 - w_j)n_t \le p_j + n_t x_j - 1$$
 $\forall j \in J$ (3.47)

Model 3.4.1: MIP model in batch-by-batch branch-and-bound

define the value of $L_{\text{max,cumul}}$, the maximum lateness of any job in the non-batched set.

Constraints (3.43) force batched jobs to start at t = 0, while (3.44) force *all* jobs to start either at t = 0 or after the last batched job ends.

Constraints (3.45) enforce a dominance rule: jobs must be assigned to the batch such that the remaining capacity, $b_r = b - \sum_j s_j x_j$, is less than the size s_j of the *smallest* job from the set of non-batched jobs that are *not longer* than the current batch. That is, if there exists an non-batched job j with $p_j \leq P_k$ and $s_j \leq b_r$, then the current assignment of jobs is

```
x_j is 1 iff job j is assigned to the batch u_{jt} is 1 iff job j starts in time slot t

T_{jt} is the set of all time slots occupied by job j if it ended at time t, that is T_{jt} = \{t - p_j + 1, \dots, t\} is the start time of the batch at the given node in the search tree L_{\max,\text{incmb}} is the incumbent (known best) value of and thus an upper bound on L_{\max} is the set of all indexed time points \{0, \dots, n_t - 1\} is 1 iff job j is either longer than the batch (p_j > P_k) or already part of the batch (x_j = 1). Neither condition must be fulfilled for constraint (3.45) to have an effect on job j
```

Table 3.1: Notation used in the decomposition model

infeasible in the model. The reasoning goes as follows: given any feasible schedule, assume there is a batch k_a with b_r remaining capacity and a later batch k_b containing a job j such that $s_j \leq b_r$ and $p_j \leq P_{k_a}$. Then job j can always be moved to batch k_a without negatively affecting the quality of the solution: if the schedule was optimal, then moving j will not affect L_{\max} at all (otherwise, it was no optimal schedule); if the schedule was not optimal, then L_{\max} will stay constant (unless j was the longest job in k_b and L_{\max} occurred in or in a batch after k_b , in which case L_{\max} will be improved).

This rule is implemented by means of a binary variable w_j , which, as defined by constraints (3.45) and (3.46), is 1 iff $p_j > P_k \lor x_j = 1$. These are the cases in which a job j is not to be considered in (3.44), and so w_j is used to scale s_j to a value insignificantly large in the eyes of the constraint's less-than relation.

After a solution is found, a child node in the search tree has run the subtree and returned, a constraint of the form

$$\sum_{j} x_{j} + \sum_{i} (1 - x_{i}) \le n_{j} - 1 \quad \forall \{j \in J | x_{j} = 1\}, \forall \{i \in J | x_{i} = 0\}$$
 (3.48)

is added to the model before the solver is called again, to exclude the last solution from the set of feasible solutions.

Using CP and cumulative packing after the batch

This approach is equivalent, but we now use CP to select the batch assignments, again based on a minimized L_{max} among the non-batched jobs. Model 3.4.2 uses interval variables j to represent the jobs; time constraints on the jobs ("est", "lft" etc.) are represented as functions such as startOf(j) and endOf(j).

Contraints (3.50) through (3.53) bi-directionally define the relationship between a job's x_j and start0f(j): $x_j = 1$ is equivalent with a start time of t = 0, and $x_j = 0$ is equivalent with a start time of $t \ge P_k$. The term $\sum_{i \in J} p_i$ is used as a large constant as it is greater than any job's start time.

Constraint (3.57) is equivalent to constraints (3.45) through (3.47) in Model 3.4.1 above. The cumulative constraint (3.60) ensures that jobs do not overlap.

(3.49)

(3.60)

s.t.
$$\operatorname{startof}(j) \geqslant (1-x_j)P_k$$
 $\forall j \in J$ (3.50) $\operatorname{startof}(j) \leqslant (1-x_j)\sum_{i \in J} p_i$ $\forall j \in J$ (3.51)
$$x_j \geqslant \frac{\frac{1}{2} - \operatorname{startof}(j)}{\sum_{i \in J} p_i} \qquad \forall j \in J \qquad \text{(3.52)}$$

$$x_j \leqslant 1 + \frac{\frac{1}{2} - \operatorname{startof}(j)}{\sum_{i \in J} p_i} \qquad \forall j \in J \qquad \text{(3.53)}$$

$$P_k \geqslant \max_{j \in J}(x_j p_j) \qquad \qquad \text{(3.54)}$$

$$\operatorname{endof}(j) = d_j + L_{\max, \text{incmb}} - 1 \qquad \forall j \in J \qquad \text{(3.55)}$$

$$L_{\max} \geqslant \max_{j \in J}(\operatorname{endof}(j) - d_j) \qquad \qquad \text{(3.56)}$$

$$\operatorname{IfThen}(p_j \leqslant P_k \land x_j = 0, b - \sum_{j \in J} s_j x_j \leqslant s_j) \quad \forall j \in J \qquad \text{(3.57)}$$

$$P_k + \frac{1}{b} \sum_i s_i p_i \leqslant d_j + L_{\max, \text{incmb}} - 1 - v_k \qquad \forall j \in J, \forall \{i \in J | d_i \leqslant d_j\} \qquad \text{(3.58)}$$

$$\sum_i x_j \geqslant 1 \qquad \qquad \forall \{j \in J | d_j = \min(d_j)\} \qquad \text{(3.59)}$$

Min.

cumul(J, b)

Model 3.4.2: CP model in batch-by-batch branch-and-bound

Results

4.1 Empirical comparison of described models

The models were tested on a set of job lists. Malapert's paper uses benchmark job lists by Daste et al. [2008a,b]. Since neither publication is available online, I created my own set of randomized job lists for the purposes of this paper, with s_j , $p_j \in [1,20]$ and $d_j \in [1,10n_j]$ where n_j is the number of jobs. Ten different sample job sets are used per unique value of n_j . The times shown in figure 4.1 are averaged over those ten instances for each n_j .

The models were run on an i7 Q740 CPU in single-thread mode, with 8 GB RAM. Solving was aborted after a time of 3600 seconds (1 hour).

The CP branch-and-bound model times out on most instances and is not shown here. The CP model times out on one 12-job instance and gets progressively worse with more jobs; similar to Malapert's original MIP model.

The final version of this document will include all results for 40 instances per n_j and for all models in tabulated form, as well as graphs to show how average solution quality (LB/UB gap) improves over time.

¹These instances will be updated to reflect the feature distributions used in Daste et al.

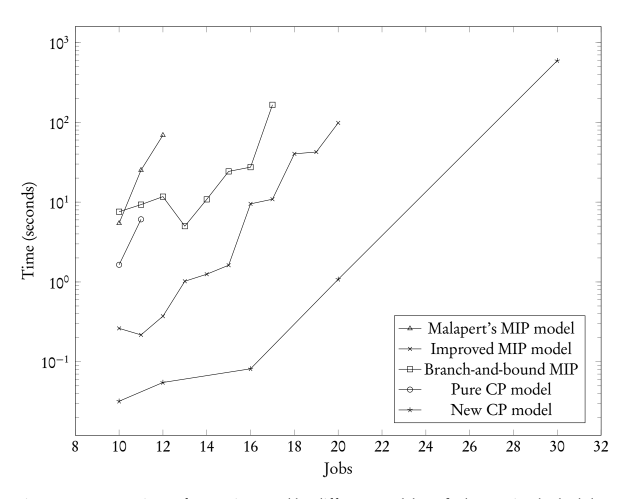


Figure 4.1: Comparison of CPU time used by different models to find an optimal schedule and prove its optimality.

Discussion

The improved MIP model currently performs best, mostly owed to constraint (3.20) which excludes a large number of potential batch assignments.

The batch-by-batch decomposition using MIP to assemble batches, while an improvement over Malapert's MIP model, is not satisfactory; the relaxations are too lax for the decomposition to compensate for the overhead of a naive branch-and-bound search.

The goal is to solve instances with up to 50 jobs in less than an hour (a performance comparable to that of Malapert's global constraint).

Future Work

6.1 Schedule

Current work on a new MIP model (not shown in this document) will continue. I will particularly try to incorporate bounds on $L_{\rm max}$ and $C_{\rm max}$ as given in papers such as [Azizoglu and Webster, 2000] and [Dupont and Dhaenens-Flipo, 2002]. Some of the CP model improvements described below shall be implemented as well, and I intend to finish the experiments by late Feburary.

The next step, an extensive discussion of the results, is planned for early March. This will involve researching how different test instances affect solving time in different models. A final draft of the thesis shall be finished by mid-March.

6.2 MIP model

6.2.1 Upper bound on L_{max}

An upper bound on $L_{\rm max}$ can be found by using a dispatch rule to find a feasible, if not optimal, schedule. A good approach could be the "best-fit" heuristic proposed in Malapert's paper.

6.2.2 Bounding the number of batches n_k

Initially, the number of batches needed is assumed to be equal to the number of jobs: $n_k = n_j$. Reducing n_k by pre-computing the maximum number of batches needed shrinks

the x_{jk} matrix, and prunes potential search branches in branch-and-bound decomposition approaches.

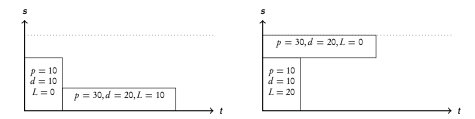


Figure 6.1: Overzealous batch elimination can increase L_{max}

Unfortunately, we cannot make a general statement that optimal solutions never have more batches than other feasible solutions – a simple counterexample is shown in figure 6.1.¹

Starting out with a one-job-per-batch schedule sorted by EDD, we can explore all feasible batch configurations recursively. To generate any other feasible schedule (including the optimal solution), jobs j are rescheduled ("moved back") from their original batch k_{origin} into a prior batch k_{earlier} . This eliminates k_{origin} and requires, of course, that k_{earlier} has sufficient capacity. If the job's processing time p_j exceeds that of k_{earlier} , then the lateness of batches between k_{earlier} and k_{origin} will increase; the merit of such a move cannot be judged a priori, so it is an *unsafe* batch elimination. *Safe* eliminations, on the other hand, will never worsen L_{max} , and only they can be considered when bounding n_k a priori.

Algorithm 2 outlines a recursive method to find an upper bound on n_k , recognizing safe batch eliminations only.

This algorithm evidently requires exponential time. A relaxed variant is a possible option: if only a single unsafe move *into* a batch is possible, no safe eliminations into that batch are considered at all and we skip to the next batch. This would greatly speed up the recursion but also significantly weaken the usefulness of the resulting upper bound.

In a branch-and-bound decomposition approach in which batches are modelled individually, an upper bound on n_k could be used to limit the depth of the search tree, or, in

¹To be more precise, we cannot state that at least one optimal solution is in the subset of feasible solutions that uses the fewest number of batches – a dominance situation that could be exploited, were it true.

Algorithm 2 Recursive algorithm to find an upper bound on n_k

combination with a method to determine a lower bound on the remaining jobs' n_k at every node, to actively prune the search tree during the search. The latter method, however, would also run in exponential time as it, again, would require knapsack-type reasoning unless we use a much less powerful relaxation.

6.3 CP Model

6.3.1 Constraint on number of batches with length $P_k > p$

Since batches take on the processing time of their longest job, there is at least one batch with $P = \max_{j} p_{j}$. We can proceed to fill batches with jobs, ordered by non-increasing processing time, based on algorithm 3.

At the end of this algorithm, we can state:

$$\texttt{globalCardinality}(\{1,\ldots,n_j\},\{P_{k,\min},\ldots,P_{k-1,\min}-1\},P_k) \quad \forall k \in \{k_0,\ldots,k_{LB(n_k)}\}, \ (6.1)$$

where the constraint takes three arguments (cards, vals and vars) and $P_{k,\min}$ denotes the minimum length of batch k.

The algorithm sorts jobs by non-increasing p, and then fills batches job by job. If a job fits into a previous batch, it is assigned there. If a job fits into multiple previous batches, it is assigned to the batch with the smallest remaining capacity. This is called *best-fit decreasing* rule, and works as follows: let J^* be the set of jobs sorted by p, then at least one batch will

Algorithm 3 Generating lower bounds on batch lengths

```
▶ initialize all jobs as unassigned jobs
n_k \leftarrow 1; S_k \leftarrow \{0\}; P_{k, \min} \leftarrow \{0\}
                                                                                       ▶ Create one empty batch of size and length zero
sort J^* by processing time, non-increasing
                                                                                               ▶ select job for assignment, longest job first
    i \leftarrow I^*.pop()
    loop through all n_k existing batches k, first batch first
                                                                                                                              ▶ no feasible batch
                                                                                       > currently known minimum remaining capacity
         if s_j < b - S_k and b - S_k < c_{\min} then
             k_p \leftarrow k_p; c_{\min} \leftarrow b - S_k
    end loop
    if |k_n| = 1 then
                                                                                                                      \triangleright assign job j to batch k_p
         S_{k_p} \leftarrow S_{k_p} + s_j
         if n_k < LB(n_k) then
              n_k \leftarrow n_k + 1
                                                                                                                                ▶ open new batch
             S_{n_k} \leftarrow s_j; P_{n_k, \min} \leftarrow p_j
                                                                                                          \triangleright assign s_i and p_i to the new batch
              leave the loop now and end.
         end if
    end if
until J^* is empty
```

be as long as the longest job j_1^* . If the next n jobs fit into this batch, then there is at least one batch not shorter than j_{n+1}^* , and similarly for subsequent batches.

Unfortunately, the optimal solution may perform better than the packing heuristic in terms of "vertical" (s_j) bin packing, and may thus require fewer batches. We therefore need to find a lower bound $LB(n_k)$ on the number of batches, and we can only guarantee the first $LB(n_k)$ of the above constraints to hold in the optimal solution. Finding a true lower bound is a two-dimensional bin packing problem, which is NP-hard. A possible but naive lower bound is j_0 , the number of jobs, ordered by decreasing s_j , that can never fit into a batch together.

6.3.2 Constraint on number of batches with length $D_k > d$

In a similar fashion, we can determine that the second batch must be due no later than the earliest-due job j_{m+1} that can *not* fit into the first batch – if we sort jobs by due date and fit the earliest m jobs into the first batch – and so on for subsequent batches. Once again, since

best-fit decreasing may not perform optimally in terms of "vertical" packing, this may not be valid for batches beyond the known $LB(n_k)$.

6.3.3 All-different constraints on P_k and D_k

Furthermore, if all jobs have different processing times, all batches will have different processing times as well: $\operatorname{alldifferent}(P_k)$. If m out of n_j jobs have different processing times, we can still enforce k_a lldifferent (P_k, m) . Some work on k_a lldifferent constraints has been done in \cite{Gain} .

Similarly, we knows that the constraint k_alldifferent(D_k , m) must be true if m out of n_i jobs have different due dates.

6.4 Decompositions, other approaches

6.4.1 Potential heuristics

Improve the initial $L_{\text{max,incumbent}}$ A better initial upper bound on L_{max} can help prune some branches of the search tree from the outset. There are several dispatch rules (or maybe other heuristics?) that could be explored to do this better.

Improve $L_{\text{max,incumbent}}$ during search It may be useful to use a heuristic like above to "complete the schedule" once a promising partial schedule has been generated. I have yet to identify situations where this is always helpful.

6.4.2 Move-back search

Another possible way to set up a branch-and-bound search for solutions works as follows: first, consider all jobs to be "single", i.e. assigned to individual batches such that $B_j = k_j \ \forall j \in J$. Compute the L_{\max} for this schedule. Then, at every level of the search tree, move some single job j into any earlier batch $k \leq k_j$, but only if that move does not violate k's capacity.

If we start with a schedule of single jobs and only allow moving single jobs into earlier batches (and any schedule can be produced by a sequence of such moves!), we maintain the EDD ordering of batches. More importantly, such moves will never shift the position of L_{max} to the right:

In any partial schedule following EDD, let k be the batch with maximum lateness $L_k = L_{\max}$. It has processing time p_k . Then the lateness of the batch before k must be $L_{k-1} \ge L_k - p_k$ as a consequence of the EDD sequencing. Any batch following k can have a lateness no greater than $L_k - 1$.

- Moving any single job from a batch following k into a batch before k will worsen L_{\max} , but not change its position. Such a move is never necessary to arrive at an optimal solution.
- Moving back any single job from a batch before k safely² will improve L_{max} , but not change its position.
- Moving back a single pre-k job j from a batch β into an earlier batch α unsafely will reduce L_k by p_{α} ; L_{\max} may still be in k, or it may be found in any batch between (and including) α and β , since their lateness $L_{[\alpha,\dots,\beta]}$ is now increased by $p_j p_{\alpha}$.
- If the max-lateness job j is single itself, it can be moved from k into an earlier batch α . If this is done *safely*, the batch immediately preceding k will still have a lateness $L_{k-1} \ge L_k p_j$. All batches after k will have their lateness reduced by p_j , but since their maximum lateness did not exceed $L_k 1$ before the move, it will now be at least 1 less than that of batch k-1.
- If the max-lateness job j was moved back *unsafely* into a batch α , batches between and including α and k-1 now have their lateness increased by p_j-p_α , while batches after k have their lateness decreased by p_α . Again, batch L_{k-1} would exceed the maximum lateness of batches after k.

This shows that after a sequence of operations in which single jobs are moved into

² for the meaning of "safe" and "unsafe" moves, compare section 6.2.2.

earlier batches, $L_{\rm max}$ will never shift to the right. In fact, this means that all jobs after the $L_{\rm max}$ -job in a single-job EDD schedule can be ignored in the scheduling problem entirely, although this turns out to be quite inconsequential as that job is often at or near the end of the single-job EDD schedule, resulting from the fact that $L_{k-1} \geqslant L_k - p_k$. By the same token, high-quality solutions often have their $L_{\rm max}$ in an early batch. This may give rise to some sort of decomposition method in which jobs are strategically moved back, and where we are trying to move $L_{\rm max}$ to the left as much as possible.

6.5 MIP model II

$$\begin{array}{lll} \text{Min.} & L_{\max} & (6.2) \\ \text{s.t.} & \sum_{k \in K} x_{jk} = 1 & \forall j \in J & (6.3) \\ & \sum_{j \in J} s_j x_{jk} \leqslant b & \forall k \in K & (6.4) \\ & x_{jk} = 0 & \forall \{j \in J, k \in K | j \leqslant k \vee s_j + s_k > b \\ & (6.5) & \\ & L_{\max} \geqslant \ell_k + \sum_{b=0}^{k-1} \left[P_b - p_b \left(1 + \sum_{i \in K} x_{bi} \right) \right] - \text{UB}(L_{\max}) \sum_{i \in K} x_{ki} & \forall k \in K & (6.6) \\ & p_j x_{jk} \leqslant P_k & \forall j \in J, \forall k \in K \\ & (6.7) \\ & (6.8) & \end{array}$$

Model 6.5.1: Malapert's original MIP model

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Appendix: Tables

A.1 Hello. This is the first part of the first Appendix Blabla.

Appendix: Code